



Title	Csi1p recruits alp7p/TACC to the spindle pole bodies for bipolar spindle formation
Author(s)	ZHENG, F; LI, T; Jin, D; Syrovatkinac, V; Scheffler, K; Tran, P; Fu, C
Citation	Molecular Biology of the Cell, 2014, v. 25, p. 2750-2760
Issued Date	2014
URL	http://hdl.handle.net/10722/205901
Rights	Molecular Biology of the Cell. Copyright © American Society for Cell Biology.

Csi1p recruits alp7p/TACC to the spindle pole bodies for bipolar spindle formation

Fan Zheng^{a,b}, Tianpeng Li^{a,b}, Dong-yan Jin^a, Viktoriya Syrovatkina^c, Kathleen Scheffler^d,
Phong T. Tran^{c,d}, and Chuanhai Fu^{a,b}

^aDepartment of Biochemistry, University of Hong Kong, Pokfulam, Hong Kong, China; ^bHKU-Shenzhen Institute of Research and Innovation, University of Hong Kong, Shenzhen, China; ^cCell and Developmental Biology, University of Pennsylvania, Philadelphia, PA 10104; ^dInstitut Curie, Centre National de la Recherche Scientifique, Paris 75005, France

ABSTRACT Accurate chromosome segregation requires timely bipolar spindle formation during mitosis. The transforming acidic coiled-coil (TACC) family proteins and the ch-TOG family proteins are key players in bipolar spindle formation. They form a complex to stabilize spindle microtubules, mainly dependent on their localization to the centrosome (the spindle pole body [SPB] in yeast). The molecular mechanism underlying the targeting of the TACC–ch-TOG complex to the centrosome remains unclear. Here we show that the fission yeast *Schizosaccharomyces pombe* TACC orthologue alp7p is recruited to the SPB by csi1p. The csi1p-interacting region lies within the conserved TACC domain of alp7p, and the carboxyl-terminal domain of csi1p is responsible for interacting with alp7p. Compromised interaction between csi1p and alp7p impairs the localization of alp7p to the SPB during mitosis, thus delaying bipolar spindle formation and leading to anaphase B lagging chromosomes. Hence our study establishes that csi1p serves as a linking molecule tethering spindle-stabilizing factors to the SPB for promoting bipolar spindle assembly.

Monitoring Editor

Kerry S. Bloom
University of North Carolina

Received: Mar 4, 2014

Revised: Jun 20, 2014

Accepted: Jul 7, 2014

INTRODUCTION

Timely bipolar spindle assembly facilitates proper kinetochore biorientation, thereby ensuring accurate chromosome segregation during mitosis (Walczak and Heald, 2008; Tanenbaum and Medema, 2010; Silkworth and Cimini, 2012). Spindle assembly takes place in prophase, with microtubule minus ends anchored at the spindle poles and microtubule plus ends interdigitating at the spindle midzone to form an antiparallel microtubule array. Spindle

bipolarity is well established by metaphase as the interdigitating microtubules slide apart and the opposing forces within the spindle are balanced (Syrovatkina et al., 2013). Bipolar spindle formation requires synergistic coordination of microtubule-associated proteins (MAPs) and kinesin motors (Tanenbaum and Medema, 2010). In general, MAPs help organize spindle microtubules into an antiparallel microtubule array, and kinesins produce forces to elongate the array to separate two spindle poles (Fu et al., 2009; Peterman and Scholey, 2009; Syrovatkina et al., 2013). Intensive studies on bipolar spindle formation have focused on the conserved MAP PRC1/ase1p and the kinesin-5 Eg5/cut7p. PRC1/ase1p becomes more static upon cross-linking two antiparallel microtubules (Janson et al., 2007; Subramanian et al., 2013), whereas Eg5/cut7p is rotationally flexible (Kapitein et al., 2005). These structural features enable PRC1/ase1p and Eg5p/cut7 to function efficiently to maintain antiparallel spindle microtubules after spindle bipolarity has been established but nonefficiently sort near parallel spindle microtubules into an antiparallel microtubule array during early prophase. This implies that initial spindle microtubule organization may require another molecular mechanism.

One possible mechanism may involve the transforming acidic coiled-coil (TACC) family proteins, as cells lacking TACC proteins typically display abnormalities in initial bipolar spindle assembly

This article was published online ahead of print in MBoc in Press (<http://www.molbiolcell.org/cgi/doi/10.1091/mbc.E14-03-0786>) on July 23, 2014.

Author contributions: F.Z., T.L., P.T.T., and C.F. designed and carried out experiments and analyzed data. F.Z., P.T.T., and C.F. wrote the article. All authors made comments.

Address correspondence to: Phong T. Tran (P.T.T. tranp@mail.med.upenn.edu), Chuanhai Fu (chuanhai@hku.hk).

Abbreviations used: EMM, Edinburgh minimal medium; GST, glutathione S-transferase; MAP, microtubule-associated protein; MBC, methyl benzimidazole-2-yl-carbamate; SPB, spindle pole body; TACC, transforming acidic coiled-coil; YE5S, yeast extract medium supplemented with adenine, leucine, uracil, histidine, and lysine.

© 2014 Zheng et al. This article is distributed by The American Society for Cell Biology under license from the author(s). Two months after publication it is available to the public under an Attribution–Noncommercial–Share Alike 3.0 Unported Creative Commons License (<http://creativecommons.org/licenses/by-nc-sa/3.0>).

“ASCB®,” “The American Society for Cell Biology®,” and “Molecular Biology of the Cell®” are registered trademarks of The American Society of Cell Biology.

Supplemental Material can be found at:
<http://www.molbiolcell.org/content/suppl/2014/07/19/mbc.E14-03-0786.DC1.html>

(Peset and Vernos, 2008). Members of this family are characterized by the conserved TACC domain at their carboxyl termini and share a similar intracellular localization pattern and conserved functions in a wide range of organisms, including yeast (Sato *et al.*, 2004), *Caenorhabditis elegans* (Bellanger and Gonczy, 2003; Srayko *et al.*, 2003), *Drosophila* (Gergely *et al.*, 2000b), and mammals (Gergely *et al.*, 2003). During mitosis, TACC proteins mainly localize to spindle microtubules and the centrosome (the spindle pole body [SPB]) in yeast; Sato *et al.*, 2004; Peset and Vernos, 2008), where they form a complex with the conserved microtubule polymerase ch-TOG proteins via their TACC domains to stabilize kinetochore microtubules (Royle, 2012) and promote spindle formation (Peset and Vernos, 2008).

Clathrin, a key protein involved in membrane trafficking, has been reported to interact with TACC3 and subsequently form the clathrin–ch-TOG–TACC3 complex for localization to spindle microtubules (Fu *et al.*, 2010; Lin *et al.*, 2010; Booth *et al.*, 2011; Hood *et al.*, 2013), resulting in kinetochore fiber stabilization, an important process for spindle stabilization (Booth *et al.*, 2011). Despite this, it remains unclear how TACC proteins are recruited to the centrosome/SPB, an equally important process required for spindle assembly and stabilization. Intriguingly, the conserved TACC domain is necessary for the localization of TACC proteins to the centrosome/SPB, independent of microtubules (Bellanger and Gonczy, 2003) and ch-TOG proteins (Gergely *et al.*, 2000b; Sato *et al.*, 2004), and the TACC domain does not interact with clathrin (Hood *et al.*, 2013). These findings highlight the importance of the TACC domain in interacting with a centrosomal/SPB protein that remains to be determined.

The fission yeast centrosome/SPB protein *csi1p* is emerging as a key protein for ensuring faithful mitotic chromosome segregation (Hou *et al.*, 2012). It is recruited to the SPB by the conserved SUN-domain protein *sad1p* (SUN1 in humans) for centromere clustering during interphase (Hou *et al.*, 2012). *Csi1p* has also been implicated in spindle formation (Costa *et al.*, 2013).

In this study, we show that *csi1p* promotes bipolar spindle assembly by recruiting the TACC orthologue *alp7p* to the SPB. The interaction domains in *csi1p* and *alp7p* lie at their carboxyl termini. When the interaction between *csi1p* and *alp7p* is compromised, *alp7p* and its binding protein *alp14p*/ch-TOG are absent from the SPBs, leading to transient monopolar spindle formation and subsequent anaphase B lagging chromosomes. Thus this work defines a new molecular mechanism regulating the SPB localization of the *alp7p*/TACC-*alp14p*/ch-TOG complex and highlights the importance of this SPB complex in bipolar spindle formation and faithful chromosome segregation.

RESULTS

Csi1p interacts with *alp7p*

Csi1p has been shown to be an essential protein for centromere clustering during interphase (Hou *et al.*, 2012). Our live-cell imaging screen identified *csi1p* as a key component for bipolar spindle formation (Costa *et al.*, 2013; see later discussion of Figure 2A). To understand the role of *csi1p* in bipolar spindle formation, we carried out a yeast-two hybrid screen for *csi1p*-binding proteins. In this screen, an *alp7p* mutant lacking the last 31 residues at its carboxyl terminus was identified as a strong interacting protein of *csi1p* (Figure 1A), and the interaction between *csi1p* and *alp7p* was further confirmed by glutathione S-transferase (GST) pull-down assays, using full-length recombinant proteins histidine (His)-*alp7p* and GST-*csi1p* (Figure 1B). Therefore the carboxyl-terminal 31 residues of *alp7p* are not required for its interaction with *csi1p*, although these residues appear to affect the *alp7p*-*csi1p* interaction specifi-

cally in budding yeast in the yeast two-hybrid assays (see later discussion of Figure 5A).

We then used live-cell imaging to examine the colocalization of *alp7p*-3GFP (green fluorescent protein) and *csi1p*-TagRFP in wild-type cells. As shown in Figure 1C, *alp7p* displayed spindle localization, with strong preferential SPB localization, and colocalization with *csi1p* throughout mitosis (also see Supplemental Figure S1A). In addition, we observed *alp7p* signals between the two SPBs on the spindle, which may represent its kinetochore localization, as previously reported (Sato *et al.*, 2004; Tang *et al.*, 2013). These additional *alp7p* signals likely did not colocalize with *csi1p*, as even over-expressed *csi1p* from the *nmt1* promoter did not localize to kinetochores (Supplemental Figure S1B). Intriguingly, *csi1Δ* and *alp7Δ* double-deletion mutant displayed additive sensitivity to the microtubule-depolymerizing drug methyl benzimidazole-2-yl-carbamate (MBC) in a dose-dependent manner (Figure 1D). Taken together, these findings suggest that *csi1p* physically and genetically interacts with *alp7p*.

Csi1p and *alp7p* are required for bipolar spindle assembly

Both *csi1p* and *alp7p* are involved in regulating chromosome segregation (Supplemental Figure S2, A and B; Sato *et al.*, 2004; Hou *et al.*, 2012; Tang *et al.*, 2013). Of importance, *alp7p* is required for microtubule organization (Thadani *et al.*, 2009), mitotic spindle assembly (Sato *et al.*, 2004; Sato and Toda, 2007), and proper attachment of spindle microtubules to kinetochores (Tang *et al.*, 2013). Therefore, to assess the biological significance of the interaction between *csi1p* and *alp7p*, we first analyzed spindle dynamics, an important process for proper chromosome segregation. Live-cell imaging revealed that similar to *alp7Δ*, *csi1Δ* cells displayed defective spindle dynamics during early mitosis, with the spindles initially forming a transient monopolar structure (Figure 2, A–D, and Supplemental Figure S2, E–G). In contrast, the wild-type spindles initially emerged as a dense, dot-like structure and quickly elongated to establish a bar-like bipolar structure 1 μ m in length within 2–3 min (Figure 2, A and E). Further measurements of spindle dynamics in *csi1Δ* and *alp7Δ* mutant cells showed that a significantly longer time was required for these mutants to assemble a bipolar spindle of 1 μ m in length compared with wild-type cells (7.0 ± 2.2 and 7.2 ± 3.9 min for *csi1Δ* and *alp7Δ* cells, respectively, vs. 3.0 ± 0.8 min for wild-type cells; Figure 2, D and E).

We then analyzed centromere clustering, another key process for proper chromosome segregation. In agreement with a previous report (Hou *et al.*, 2012), 56% of the *csi1Δ* cells displayed centromere declustering (Supplemental Figure S1, C and D). However, no wild-type cells and only 2% of the *alp7Δ* cells displayed centromere declustering (Supplemental Figure S1, C and D). The high degree of similarity of spindle defect, not centromere declustering, caused by the absence of *csi1p* or *alp7p* suggests that the interaction between *csi1p* and *alp7p* is likely involved in bipolar spindle formation.

Csi1p recruits *alp7p* to the SPB during mitosis

Next we analyzed the localization interdependence of *alp7p* and *csi1p* by confocal microscopy. Fluorescence intensity analysis of sum projection images showed that the majority of *alp7p* colocalized with *csi1p* at the SPBs in wild-type cells in early mitosis, whereas most *alp7p* signals appeared between the two SPBs as several distinct dots in *csi1Δ* cells (Figure 3A). Further fluorescence intensity analysis revealed that the *alp7p* dots between the two SPBs in *csi1Δ* cells colocalized with *cnp3p* (a kinetochore protein; CENP-C in humans) at the kinetochores (Figure 3B). To exclude the effect of spindle microtubules on *alp7p* localization, we then took advantage

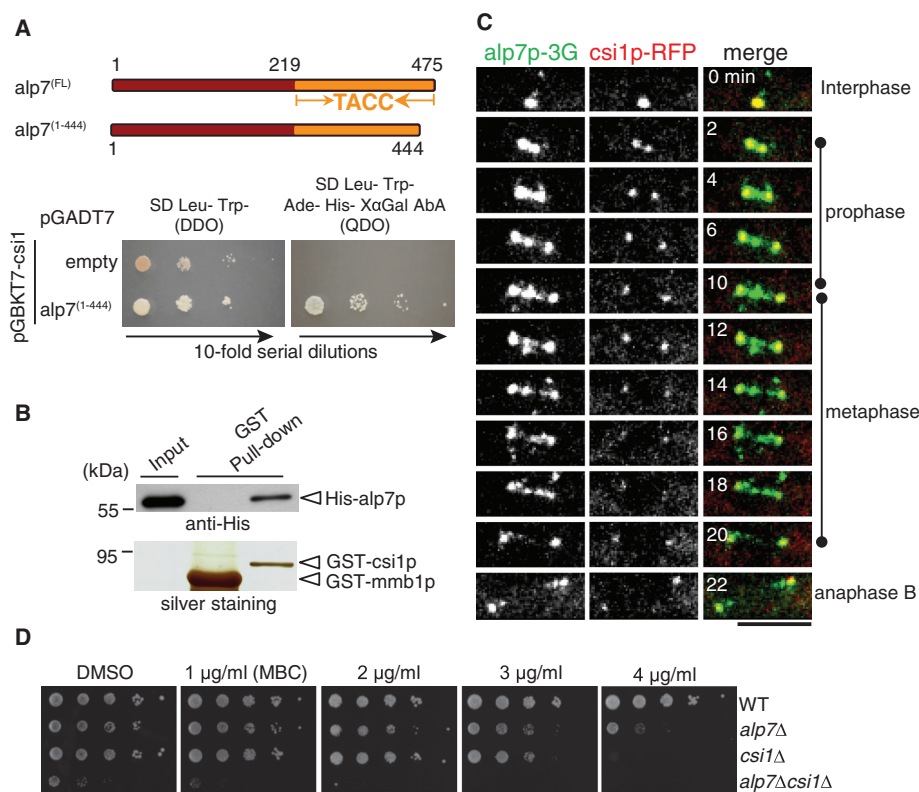


FIGURE 1: Csi1p interacts with alp7p. (A) Yeast two-hybrid assays testing the interaction between csi1p and alp7p. Y2HGold budding yeasts cotransformed with BD-csi1 and AD-alp7⁽¹⁻⁴⁴⁴⁾ or empty AD plasmids were subjected to 10-fold serial dilutions and spotted on SD/-Leu/-Trp (DDO) and SD/-Leu/-Trp/-Ade/-His (QDO) plus X- α -gal and Aureobasidin A (AbA) plates and incubated at 30°C for 4 d. (B) GST pull-down assays. Full-length recombinant proteins GST-csi1p and His-alp7p were produced in *E. coli*; the precipitation products were analyzed by Western blotting with anti-His antibody. Note that His-alp7p coprecipitated with GST-csi1p, not the control GST-mmb1p. (C) Maximum projection live-cell images of a cell expressing csi1p-TagRFP and alp7p-3GFP from their own promoters. Alp7p colocalized with csi1p at the SPBs throughout mitosis and appeared as distinct dots between the two SPBs during metaphase. Scale bar, 5 μ m. (D) MBC sensitivity assays. Tenfold serial dilutions of wild-type (WT), *alp7* Δ (*alp7p* null), *csi1* Δ (*csi1p* null), and *alp7* Δ *csi1* Δ (*alp7p* and *csi1p* null) cells were grown at 30°C for 4 d on YE5S plates containing dimethyl sulfoxide or the indicated concentrations of MBC. Note that cells lacking both *alp7*⁺ and *csi1*⁺ displayed an additive defect in cell growth.

of the *nda3*-KM311 β -tubulin thermosensitive mutant strain (Hiraoka *et al.*, 1984), which can be arrested in prophase by temperature shift to the restrictive temperature of 16°C and has no spindle upon arrest. Similarly, we observed that at the restrictive temperature, the absence of csi1p caused delocalization of alp7p from the SPBs in prophase (Figure 3C), and the delocalized alp7p colocalized with *cnp3p* at the kinetochores (Figure 3D). Consistently, the absence of csi1p also caused delocalization of the alp7p-binding partner alp14p from the SPBs (Supplemental Figure S3). Moreover, we tested conversely whether alp7p is required for the SPB localization of csi1p. The fluorescence intensity analysis showed that the SPB localization of csi1p was not altered in the absence of alp7p (Figure 3E). Therefore we conclude that csi1p is required for the SPB localization of the alp7p-alp14p complex.

The carboxyl terminus of csi1p is responsible for interacting with alp7p

In addition to centromere clustering, csi1p plays a role in bipolar spindle formation. It is likely that csi1p recruits alp7p to the SPBs for

promoting bipolar spindle formation. To assess precisely the function of the csi1p-alp7p interaction, we sought to create a separation-of-function mutant for csi1p, whose interaction with alp7p is specifically inhibited. For this purpose, we used yeast two-hybrid assays to map the key csi1p residues responsible for interacting with alp7p. Multiple attempts revealed that Ile-463 and Pro-464 within the minimal domain (amino acids [aa] 461–480) lying at the carboxyl terminus of csi1p were indispensable for interacting with alp7p (Figure 4A).

Csi1p is a low-abundance nuclear protein, with ~500 molecules in a cell (Marguerat *et al.*, 2012). Moreover, csi1p likely interacts with alp7p within a very short time window at prophase. These make it technically challenging to perform coimmunoprecipitation to test csi1p-alp7p interaction. Instead, we used the bimolecular fluorescence complementation (BiFC) assay, which has emerged as a key assay for examining protein–protein interaction in many model organisms, including yeast (Kodama and Hu, 2012). Generally, two complement peptide fragments of GFP (VN and VC in our study) are fused with proteins to be tested, and the interaction of the two fusion proteins can then bring the two complementary peptide fragments together to form a mature GFP molecule to give fluorescence signals. Consistently, BiFC assays confirmed that csi1p interacts with alp7p at the SPB (Figure 4B) and that the substitution of Ile-463 and Pro-464 for two asparagines in csi1p (designated as csi1p^{463IPNN}) can effectively inhibit its interaction with alp7p in vivo (Figure 4B). Csi1p^{463IPNN} localized to the SPB as wild-type csi1p (designated as csi1p^{WT}; Supplemental Figure S4A), and their expression levels were comparable (Figure 4H). Further,

csi1p^{463IPNN} did not cause centromere declustering (Supplemental Figure S4, B and C). However, csi1p^{463IPNN} caused delocalization of alp7p from the SPBs (Figure 4C), thus phenocopying the effect of the absence of csi1p on alp7p localization. Consistently, the delocalized alp7p colocalized with *cnp3p* at the kinetochores (Supplemental Figure S4E).

We then examined spindle dynamics in *csi1* Δ cells expressing csi1p^{463IPNN} or csi1p^{WT}. As expected, csi1p^{WT} restored normal spindle dynamics, whereas csi1p^{463IPNN} did not (Figure 4D). Further quantitative measurements also confirmed that csi1p^{463IPNN} caused transient monopolar spindle formation (Figure 4E and Supplemental Figure S4D), and significantly longer time was required for csi1p^{463IPNN} mutant cells to assemble a spindle 1 μ m in length (i.e., 7.7 ± 2.6 and 2.4 ± 0.01 min for csi1p^{463IPNN} and csi1p^{WT} cells, respectively; Figure 4F). Moreover, similar to *csi1* Δ , csi1p^{463IPNN} cells were sensitive to MBC in a dose-dependent manner (Figure 4G). Hence csi1p^{463IPNN} phenocopies the *csi1* Δ spindle defect, supporting the conclusion that csi1p recruits alp7p to the SPBs specifically for promoting bipolar spindle formation.

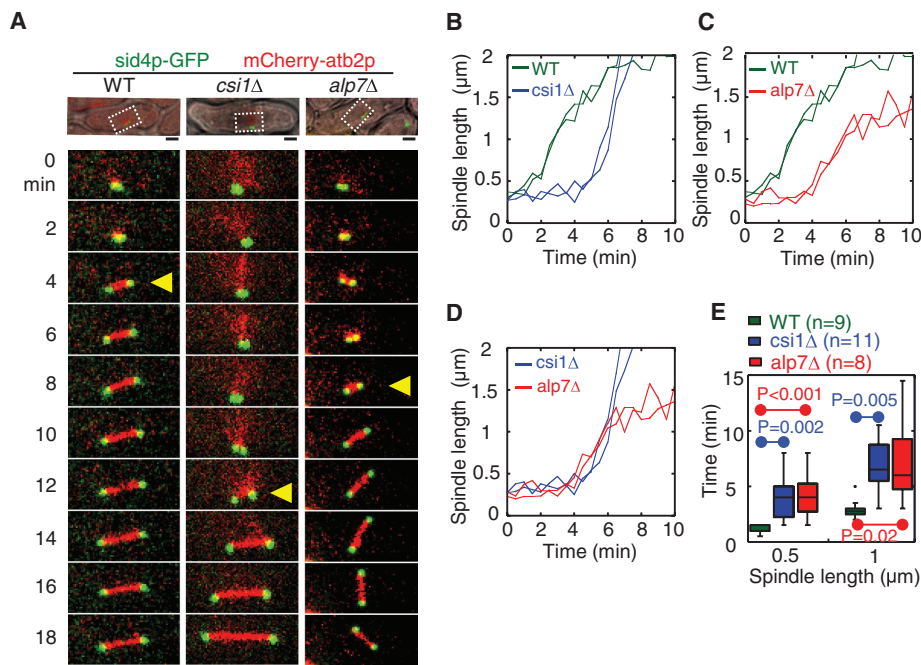


FIGURE 2: Csi1p is required for bipolar spindle assembly. (A) Maximum projection live-cell images of wild-type, *csi1Δ*, and *alp7Δ* cells expressing sid4p-GFP (SPB marker) and mCherry-atb2p (α -tubulin). Yellow triangles mark bipolar spindles $\sim 1 \mu\text{m}$ in length. Highlighted regions in the differential interference contrast pictures. Scale bar, $2 \mu\text{m}$. (B–D) Representative plots of spindle length against time for wild-type and *csi1Δ* cells (B), wild-type and *alp7Δ* cells (C), and *csi1Δ* and *alp7Δ* cells (D). (E) Box plots of time for assembly of bipolar spindles measuring 0.5 and $1 \mu\text{m}$ in length in wild-type, *csi1Δ*, and *alp7Δ* cells. Student's *t* test was used to calculate *p* values. Cell numbers analyzed are indicated.

The alp7p TACC domain is responsible for interacting with csi1p

The TACC domain is necessary and sufficient for localizing TACC proteins to the centrosome/SPB (Gergely *et al.*, 2000b; Bellanger and Gonczy, 2003; Sato *et al.*, 2004). This prompted us to explore whether the TACC domain in alp7p is responsible for interacting with csi1p. Similarly, yeast two-hybrid assays were used to map the minimal csi1p-interacting domain in alp7p. Because the last 31 residues in alp7p appeared to affect its interaction with csi1p in budding yeast (Figure 5A; full-length alp7p displayed no interaction with csi1p), we chose to use a series of alp7p deletion truncation mutants lacking the 31 residues for the yeast two-hybrid assays. This attempt revealed that residues 307–312 within the TACC domain in alp7p were required for interacting with csi1p (Figure 5A). Further, alp7p lacking residues 307–312 (designated alp7p^(Δ307–312)) showed no BiFC signals when paired with csi1p in the BiFC assays (Figure 5B), suggesting that alp7p residues 307–312 are also important for interacting with csi1p *in vivo*. We then tested the expression levels of alp7p^{WT} and alp7p^(Δ307–312) *in vivo*, showing that their expression levels were comparable (Figure 5H). Next live-cell imaging showed that alp7p^(Δ307–312) did not cause centromere declustering (Supplemental Figure S5, A and B), but its SPB localization was impaired (Figure 5C), with most alp7p^(Δ307–312) residing at the kinetochores (Supplemental Figure S5F). This led to transient monopolar spindle formation (Figure 5, D–F, and Supplemental Figure S5C). Moreover, similar to *alp7Δ*, alp7p^(Δ307–312) cells were sensitive to MBC in a dose-dependent manner at 35°C (Figure 5G). Of interest, csi1p^{463IPNN} alp7p^(Δ307–312) double-mutant cells showed no obvious additive effect on MBC sensitivity (Figure 5G), confirming that csi1p^{463IPNN} and alp7p^(Δ307–312) mutants likely operate in the same pathway.

Our further yeast two-hybrid assays showed that although the alp7p TACC domain also interacts with alp14p (Sato *et al.*, 2004; Sato and Toda, 2007), the residues 461–467 at the extreme carboxyl terminus of alp7p are what is important for interacting with alp14p (Supplemental Figure S5D). Therefore the csi1p- and alp14p-interacting regions in the alp7p TACC domain do not overlap.

Proper interaction between csi1p and alp7p is required for faithful chromosome segregation

Timely bipolar spindle formation ensures faithful chromosome segregation (Walczak and Heald, 2008; Tanenbaum and Medema, 2010; Silkworth and Cimini, 2012). We then asked whether the transient monopolar spindle formation caused by the compromised interaction between csi1p and alp7p affects faithful chromosome segregation. To address this question, we first carried out live-cell imaging to examine kinetochore dynamics in wild-type, *csi1Δ*, and *alp7Δ* cells. As shown in Figure 6A, both deletion mutants displayed remarkable anaphase B lagging chromosomes. To quantify this phenotype, we measured the percentage of anaphase B cells that displayed spindles 4 , 5 , 6 , and $7 \mu\text{m}$ in length, respectively, and concomitantly showed lagging chromosomes.

This analysis showed that a comparable large number of anaphase B *csi1Δ* and *alp7Δ* cells displayed lagging chromosomes, whereas no anaphase B wild-type cells with a spindle length $>4 \mu\text{m}$ displayed lagging chromosomes (Figure 6B). Moreover, csi1p^{WT} restored normal chromosome segregation in *csi1Δ* cells, whereas csi1p^{463IPNN}, which cannot interact with alp7p, only partially restored normal chromosome segregation in *csi1Δ* cells (Figure 6C). Likewise, alp7p^{WT} restored normal chromosome segregation in *alp7Δ* cells (Figure 6D); however, alp7p^(Δ307–312), which cannot interact with csi1p, partially rescued lagging chromosomes in *alp7Δ* cells (Figure 6D). Further, csi1p^{463IPNN} alp7p^(Δ307–312) double-mutant cells showed a comparable degree of anaphase B chromosome lagging as single alp7p^(Δ307–312) mutant cells (Figure 6D), further confirming that csi1p^{463IPNN} and alp7p^(Δ307–312) mutants operate in the same pathway. Taken together, these findings suggest that timely bipolar spindle formation mediated by the csi1p-alp7p complex at the SPB is required for faithful chromosome segregation.

DISCUSSION

Spindle assembly and stabilization requires the Alp7p/TACC-alp14p/ch-TOG complex, which targets to kinetochore microtubules (Royle, 2012) and the spindle poles (Sato *et al.*, 2004; Peset and Vernos, 2008; Royle, 2012). Although it is relatively clear that TACC and ch-TOG interact with the trimerized clathrin heavy chains (CHCs) to cross-bridge kinetochore microtubules for spindle stabilization (Fu *et al.*, 2010; Lin *et al.*, 2010; Hood *et al.*, 2013), it remains unknown how the Alp7p/TACC-alp14p/ch-TOG complex is tethered to the spindle poles, a key process for timely bipolar spindle formation (Sato *et al.*, 2004; Peset and Vernos, 2008). Here we demonstrate that alp7p and alp14p depend on csi1p for localization to the

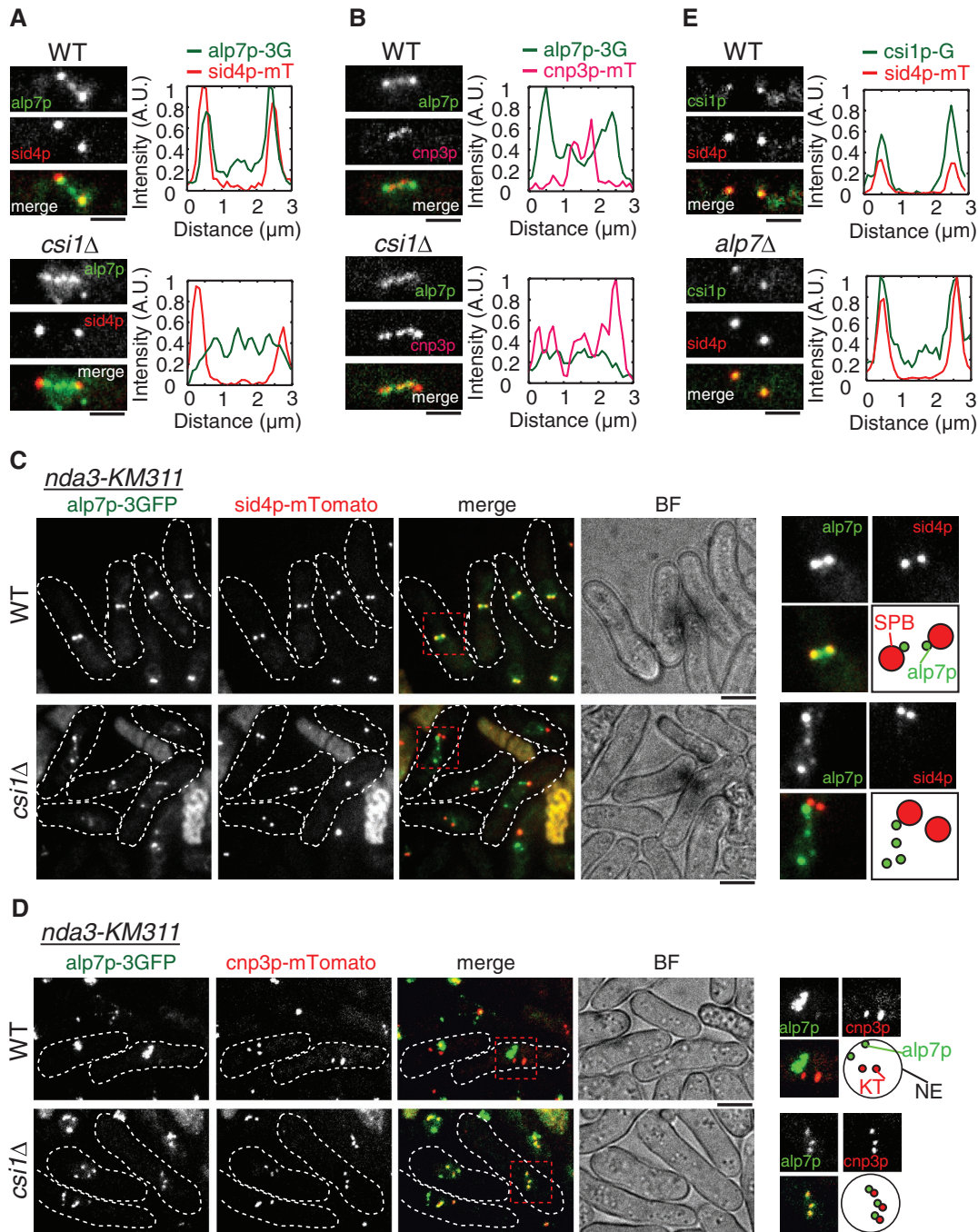


FIGURE 3: Csi1p recruits alp7p to the SPBs. (A) Sum projection images of wild-type and *csi1Δ* cells expressing alp7p-3GFP and the SPB marker sid4p-mTomato (SPB marker) from their own promoters. Fluorescence intensity measurements were carried out with MetaMorph to analyze signal profiles of alp7p and sid4p along the spindle. Alp7p colocalized with sid4p at the SPBs in the wild-type cell but not in the *csi1Δ* cell; in the *csi1Δ* cell, alp7p appeared as distinct dots between the two SPBs. Scale bars, 2 μm. (B) Sum projection images of wild-type and *csi1Δ* cells expressing alp7p-3GFP and the kinetochore marker cnp3p-mTomato (kinetochore marker) from their own promoters. Fluorescence intensity measurements were carried out to analyze signal profiles of alp7p and cnp3p along the spindle. Alp7p colocalized with cnp3p at the kinetochores in the *csi1Δ* cell but not in the wild-type cell. Scale bars, 2 μm. (C) Sum projection images of *nda3-KM311* wild-type and *nda3-KM311 csi1Δ* cells expressing alp7p-3GFP and sid4p-mTomato. Before imaging, *nda3-KM311* cells were cultured at their restrictive temperature 16°C for 6 h, a condition that can efficiently disassemble the spindles to arrest the cells at prophase/prometaphase. Magnified views on the right highlight the delocalization of alp7p from the SPBs in the absence of csi1p. Scale bars, 2 μm. (D) Sum projection images of *nda3-KM311* wild-type and *nda3-KM311 csi1Δ* cells expressing alp7p-3GFP and cnp3p-mTomato. Before imaging, *nda3-KM311* cells were cultured at their restrictive temperature 16°C for 6 h. Magnified views on the right highlight the colocalization of alp7p and cnp3p in the absence of csi1p. Scale bars, 2 μm. (E) Sum projection images of wild-type and *alp7Δ* cells expressing csi1p-GFP and sid4p-mTomato from their own promoters. Fluorescence intensity measurements were carried out to analyze signal profiles of csi1p and sid4p along the spindle. Csi1p colocalized with sid4p at the SPBs in both wild-type and *alp7Δ* cells. Scale bars, 2 μm.

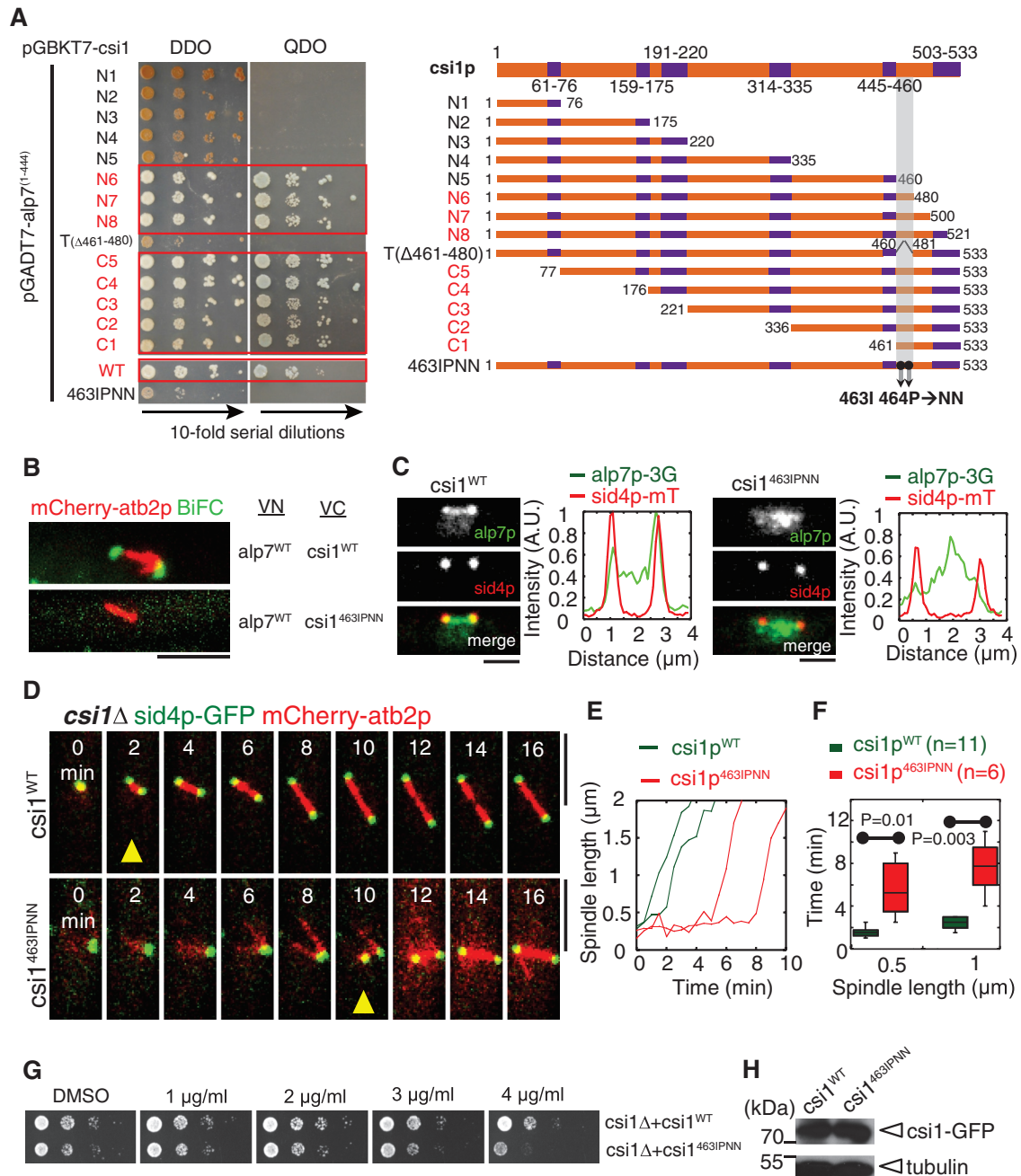


FIGURE 4: Csi1p carboxyl terminus is responsible for interacting with alp7p. (A) Yeast two-hybrid assays for mapping the minimal alp7p interaction domain in csi1p. A series of csi1p deletion truncation mutants, as indicated in the schematic diagram (coiled-coil domains indicated in purple), was used to test their interaction with alp7¹⁻⁴⁴⁴, revealing that domain 461–480 at the csi1p carboxyl terminus is important for interacting with alp7¹⁻⁴⁴⁴. Further, the two residues Ile-463 and Pro-464 within the minimal domain in csi1p are key residues responsible for the interaction with alp7p. (B) BiFC assays. Maximum projection images of cells expressing alp7^{WT}-VN and csi1^{WT}-VC or csi1^{463IPNN}-VC from the *nmt1* promoter. Cells were cultured in EMM (Edinburgh minimal medium) without thiamine for 14 h before imaging. Note that only the cell expressing wild-type csi1p gave BiFC signals. Scale bars, 5 μ m. (C) Maximum projection images of *csi1* Δ cells expressing alp7p-3GFP, sid4p-mTomato, and either wild-type csi1p (indicated as csi1^{WT}) or mutant csi1^{463IPNN} (indicated as csi1^{463IPNN}) from a csi1p promoter at the *leu1-32* locus. Fluorescence intensity measurements were carried out to analyze alp7p signal profiles along the spindles. Alp7p no longer concentrated at the SPBs in the csi1^{463IPNN} cell. Scale bars, 2 μ m. (D) Maximum projection live-cell images of csi1^{WT} and csi1^{463IPNN} cells expressing sid4p-GFP and mCherry-*atb2p*. Yellow triangles mark bipolar spindles \sim 1 μ m in length. Note that the csi1^{463IPNN} cell displayed transient monopolar spindle formation. Scale bars, 5 μ m. (E) Representative plots of spindle length against time for csi1^{WT} and csi1^{463IPNN} cells. (F) Box plots of time for assembly of bipolar spindles measuring 0.5 and 1 μ m in length in csi1^{WT} and csi1^{463IPNN} cells. Student's *t* test was used to calculate *p* values. Cell numbers analyzed are indicated. (G) MBC sensitivity assays for csi1^{WT} and csi1^{463IPNN} cells. The cells were grown at 30°C for 4 d. Similar to csi1 Δ , csi1^{463IPNN} cells were sensitive to MBC. (H) Western blot analysis of cells expressing csi1^{WT}-GFP and csi1^{463IPNN}-GFP.

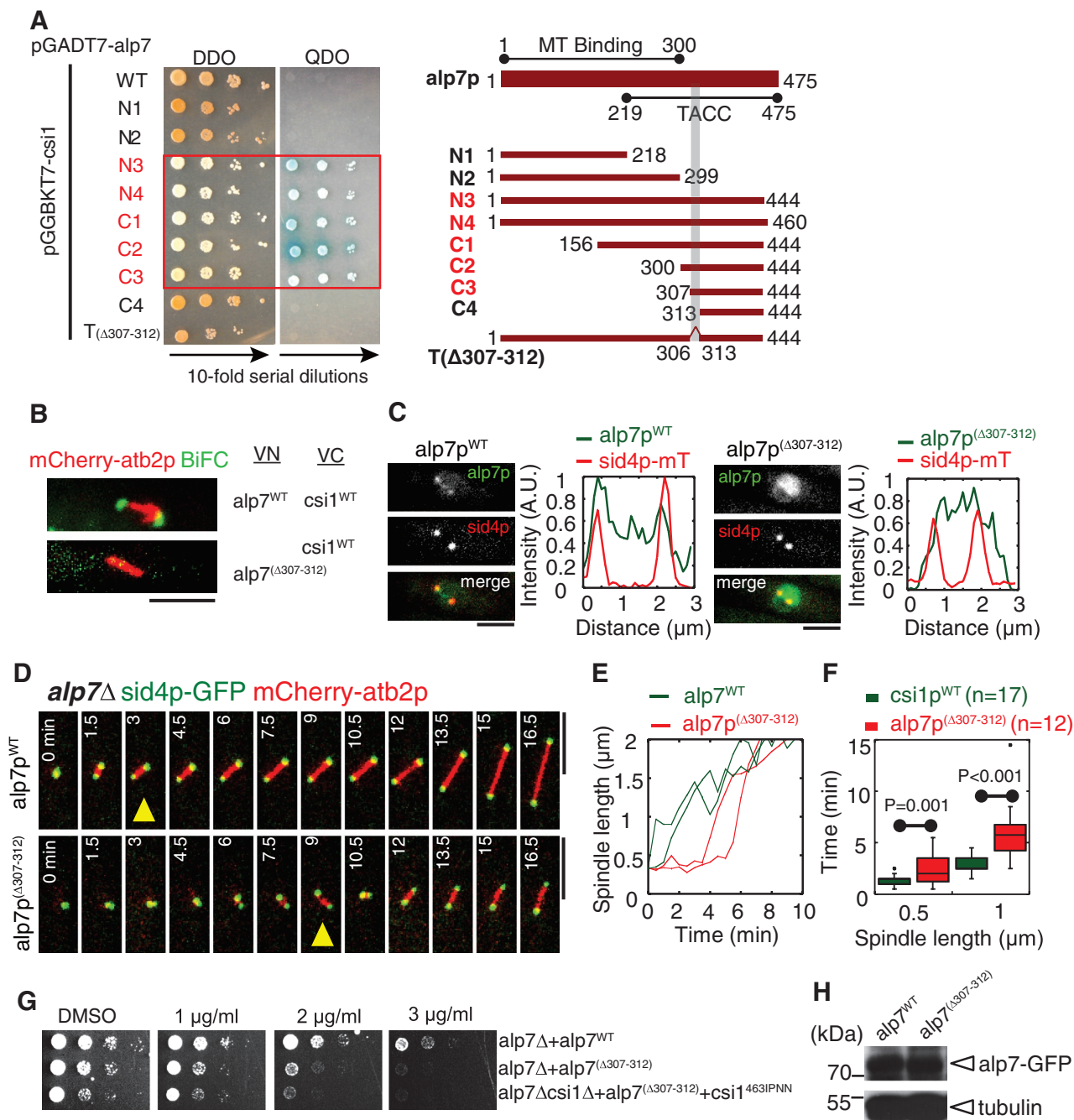


FIGURE 5: The alp7p TACC domain is responsible for interacting with csi1p. (A) Yeast two-hybrid assays for mapping the minimal csi1p interaction domain in csi1p. A series of alp7p deletion truncation mutants, as indicated in the schematic diagram, was used to test their interaction with csi1p, revealing that the domain 307–312 within the alp7p TACC domain is necessary for interacting with csi1p. (B) BiFC assays. Maximum projection images of cells expressing alp7p^{WT}-VN or alp7p(Δ 307-312)-VN and csi1p^{WT}-VC from the nmt1 promoter. Cells were cultured in EMM without thiamine for 14 h before imaging. Note that only the cell expressing wild-type alp7p gave BiFC signals. Scale bars, 5 μ m. (C) Maximum projection images of alp7 Δ cells expressing sid4p-mTomato and either wild-type alp7p (indicated as alp7p^{WT}) or mutant alp7p(Δ 307-312) (indicated as alp7p(Δ 307-312)) from a alp7p promoter at the leu1-32 locus. Fluorescence intensity measurements were carried out to analyze alp7p signal profiles along the spindles. alp7p(Δ 307-312) no longer concentrated at the SPBs. Scale bars, 2 μ m. (D) Maximum projection live-cell images of alp7p^{WT} and alp7p(Δ 307-312) cells expressing sid4p-GFP and mCherry-atb2p. Yellow triangles mark bipolar spindles ~1 μ m in length. Bipolar spindle assembly in the alp7p(Δ 307-312) was impaired. Scale bars, 5 μ m. (E) Representative plots of spindle length against time for alp7p^{WT} and alp7p(Δ 307-312) cells. (F) Box plots of time for assembly of bipolar spindles measuring 0.5 and 1 μ m in length in alp7p^{WT} and alp7p(Δ 307-312) cells. Student's t test was used to calculate p values. Cell numbers analyzed are indicated. (G) MBC sensitivity assays for alp7^{WT}, alp7(Δ 307-312), and csi1^{463IPNN} and alp7(Δ 307-312) double-mutant cells. The cells were grown at 35°C for 4 d. Similar to alp7 Δ , alp7(Δ 307-312) cells were sensitive to MBC, and csi1^{463IPNN} and alp7(Δ 307-312) displayed no obvious additive effect. (H) Western blot analysis of cells expressing alp7^{WT}-GFP and alp7(Δ 307-312)-GFP.

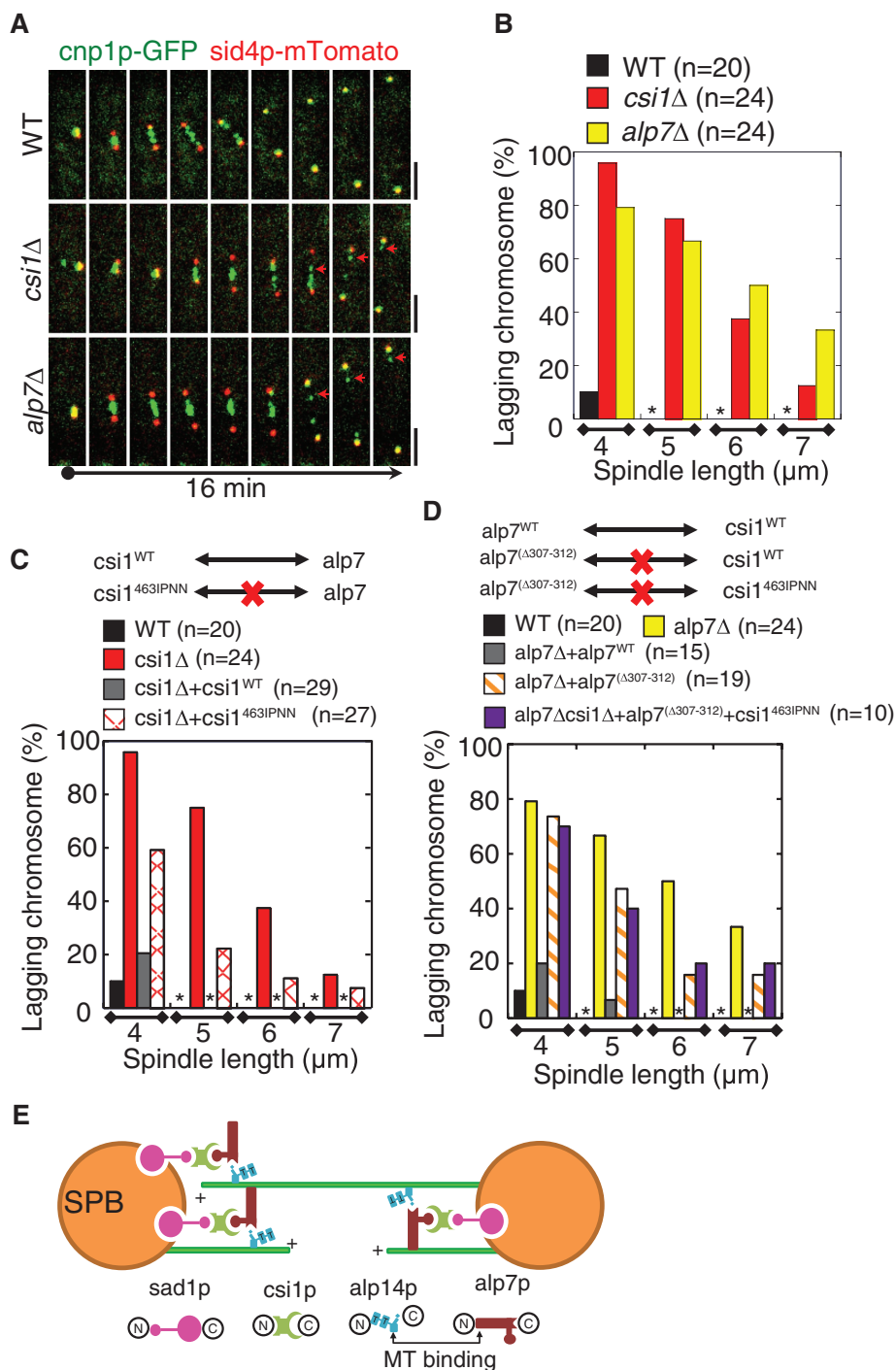


FIGURE 6: The coordination between *csi1p* and *alp7p* is required for accurate chromosome segregation. (A) Maximum projection live-cell images of wild-type, *csi1Δ*, and *alp7Δ* cells expressing *cnp1p*-GFP and *sid4p*-mTomato from their own promoters. Anaphase B lagging chromosomes were detected in *csi1Δ* and *alp7Δ* cells (arrows). Scale bars, 2 μm. (B–D) Graphs of the percentage of wild-type, *csi1Δ*, and *alp7Δ* cells (B), wild-type, *csi1Δ*, *csi1*^{WT}, and *csi1*^{463IPNN} cells (C), and wild-type, *alp7Δ*, *alp7p*^{WT}, *alp7p*^{Δ307-312}, and *csi1*^{463IPNN} and *alp7*^{Δ307-312} double-mutant cells (D) displaying anaphase B lagging chromosomes. Quantification was categorized according to the spindle length (4, 5, 6, and 7 μm, respectively). Stars indicate no anaphase B lagging chromosomes. Note that *csi1*^{463IPNN} and *alp7*^{Δ307-312} has no additive effect on anaphase B lagging chromosomes. Cell numbers analyzed are indicated. (E) A model for bipolar spindle assembly. *Csi1p* serves as a linking molecule between *sad1p* and *alp7p* to recruit the *alp7p*-*alp14p* complex to the SPBs for promoting bipolar spindle formation. The N-termini of *sad1p* and *csi1p* (Hou et al., 2012) and the C-termini of *alp7p* and *alp14p* (Sato et al., 2004) bind to each other, respectively. The microtubule-binding domains lie at the N-terminus (aa 1–300) of *alp7p*, outside of the TACC domain (Thadani et al., 2009), and the C-terminus of *alp14p*,

SPBs (Figure 3), and the SPB localization of *alp7p* and *alp14p* then promotes timely bipolar spindle formation (Figures 2, 4, and 5), thus ensuring faithful chromosome segregation (Figure 6). We propose that *csi1p* serves as a linking molecule, recruiting the microtubule-stabilizing factor *alp7p*-*alp14p* complex to the SPB, where this complex facilitates the lateral binding of adjacent microtubules for promoting bipolar spindle formation (Figure 6E).

In addition to localizing to spindle microtubules (Royle, 2012), the evolutionarily conserved *alp7p*/TACC-*alp14p*/ch-TOG complex shares another striking similarity in concentrating at the centrosomes in higher eukaryotic cells (Peset and Vernos, 2008) and the SPBs in yeast during mitosis (Usui et al., 2003; Sato et al., 2004). The targeting of the *alp7p*/TACC-*alp14p*/ch-TOG complex to the centrosome/SPB is a decisive step for mitotic spindle assembly in *Drosophila* (Gergely et al., 2000b) and *C. elegans* (Bellanger and Gonczy, 2003; Le Bot et al., 2003; Srayko et al., 2003) embryos, *Xenopus* egg extracts (O'Brien et al., 2005; Peset et al., 2005), human somatic cells (Gergely et al., 2000a, 2003), and yeasts (Usui et al., 2003; Sato et al., 2004). The mitotic kinase Aurora A has been shown to be essential for targeting TACC proteins to the centrosome (Giet et al., 2002; Bellanger and Gonczy, 2003; Le Bot et al., 2003; Peset et al., 2005). However, the sole yeast Aurora kinase, *ark1*, does not localize to the SPB (Petersen et al., 2001), and thus it is unlikely that it regulates the SPB localization of the yeast TACC *alp7p*. Moreover, *alp7p* contains no transmembrane domains, making it also unlikely for *alp7p* to localize to the SPB by inserting into the nuclear envelope. Intriguingly, the *C. elegans* TACC protein TACC-1 localizes to the centrosome independent of microtubules (Bellanger and Gonczy, 2003), and the conserved TACC domain is necessary and sufficient for localization to the centrosome in *Drosophila* embryos (Gergely et al., 2000b). In fission yeast, *alp7p*/TACC does not depend on *alp14p*/ch-TOG (Sato et al., 2004) or on microtubules (Figure 3C) for localizing to the SPB. All these findings raise the possibility that a conserved centrosomal/SPB protein or its

adjacent to the dual TOG domains (indicated as T; Al-Bassam et al., 2012), respectively. *Csi1p* recruits *alp7p* to the SPBs through an interaction between *csi1p* C-terminus (aa 461–480) and a region (aa 307–312) in the TACC domain of *alp7p* (indicated as the protruding circular shape).

functional homologues may be responsible for recruiting alp7p/TACC to the centrosome/SPB. Knowledge of such proteins is beginning to emerge. For example, NDEL1, a centrosomal protein involved in dynein function, is required for targeting TACC3 to the centrosome (Mori *et al.*, 2007), and the centrosomin Cnn is required for proper localization of D-TACC to *Drosophila* embryonic centrosomes (Zhang and Megraw, 2007).

Csi1p has no amino acid sequence similarity to either NDEL1 or Cnn. However, the absence of csi1p causes nearly identical defects of spindle formation and chromosome segregation as the absence of alp7p (Figures 2 and 6, A and B), highlighting that the two proteins may function in the same pathway. Indeed, csi1p determines the SPB localization of alp7p, but the converse is not true (Figure 3). Csi1p is also required for the SPB localization of alp14p (Supplemental Figure S3). This may be via alp7p, as there is no direct interaction between csi1p and alp14p (Supplemental Figure S5E). The alp7p-alp14p complex is a key target of the Ran GTPase-dependent spindle assembly machinery (Sato and Toda, 2007). Our present work further extends this model, in which the Ran machinery is shown to target the alp7p-alp14p complex to the nucleus for accumulation. Thus our data suggest that upon entering the nucleus at mitosis onset, the alp7p-alp14p complex is tethered to the SPBs by csi1p for promoting bipolar spindle formation (Figure 3). The absence of csi1p, that is, losing the SPB docking site for alp7p, leads to alp7p mislocalization, with the majority of alp7p localizing to the kinetochores (Figure 3, B and D), consistent with the recent finding that alp7p can also dock at the kinetochores by interacting with the internal loop of the kinetochore protein Ndc80 (Tang *et al.*, 2013). Of importance, in wild-type cells, alp7p does not display strong localization at the kinetochore in prophase and instead mainly localizes to the SPBs (Figure 3, B and C). Hence, by interacting with csi1p, alp7p is confined to the SPB region and thus is biased toward promoting bipolar spindle formation during early mitosis.

How does then the SPB localization of the alp7p-alp14p complex promote bipolar spindle formation? Csi1p is recruited to the SPBs by the SUN-domain protein sad1p through a physical interaction between their amino termini (Hou *et al.*, 2012; Figure 6E). Our domain-mapping data further show that the extreme carboxyl terminus of csi1p (Figure 4A) interacts with a small region within the TACC domain of alp7p (Figure 5A) to recruit alp7p to the SPBs (Figure 6E). This interaction likely does not affect the interaction between alp7p and alp14p, given that the alp14p-interacting domain lies at the end of the alp7p carboxy terminus (Supplemental Figure S5D), not overlapping with the csi1p-interacting domain. These unique arrangements of domain structure enable alp7p not only to be tethered to the SPBs, but also simultaneously to form a complex with the microtubule-stabilizing factor/microtubule polymerase alp14p (Al-Bassam *et al.*, 2012). Because alp7p and alp14p interact with each other through their extreme carboxyl termini (Sato *et al.*, 2004; Supplemental Figure S5D), the two microtubule-binding domains, respectively from alp7p and alp14p as previously defined (Thadani *et al.*, 2009; Al-Bassam *et al.*, 2012), flank the csi1p-binding site in the alp7p-alp14 heterodimer, a configuration favorable for bundling microtubules. The absence of the alp7p-alp14 complex at the SPB results in monopolar spindles with protruding microtubules (Figures 2, 4D, and 5C), suggesting that the alp7p-alp14p complex at the SPBs can efficiently cross-bridge adjacent microtubules that just emerge from the two SPBs during prophase and thus allows for formation of the bar-like spindle in yeast. Hence csi1p recruits to the SPBs the microtubule-bundling and -stabilizing factor alp7p-alp14p complex to promote timely bipolar spindle formation.

Thus far csi1p has been shown to play roles in centromere clustering (Hou *et al.*, 2012) and bipolar spindle formation (Figures 2 and 6E). Therefore, the lagging chromosomes caused by the absence of csi1p could be due to either centromere declustering or transient monopolar spindle formation. Given that the mutant csi1p^{463IPNN} causes transient monopolar spindle formation (Figure 4D) but not centromere declustering (Supplemental Figure S4, B and C), it is an ideal separation-of-function mutant for assessing the individual contributions of centromere clustering and bipolar spindle formation to faithful chromosome segregation. Csi1p-^{463IPNN} still causes anaphase B lagging chromosomes but to a lesser degree than csi1Δ (Figure 6C and Supplemental Figure S6A), suggesting that not only bipolar spindle formation but also centromere clustering contributes to faithful chromosome segregation. Consistently, both alp7Δ and alp7p^(Δ307-312) mutants, which display normal centromere clustering (Supplemental Figures S2, C and D, and S5, A and B) but transient monopolar spindles (Figures 2 and 5D), show massive anaphase B lagging chromosomes (Figure 6, A and D, and Supplemental Figure S6B), reinforcing the conclusion that proper bipolar spindle formation is required for faithful chromosome segregation. Thus csi1p ensures faithful chromosome segregation not only by clustering centromeres at the SPBs (Hou *et al.*, 2012), but also by promoting timely bipolar spindle formation.

Blast analysis shows no homologues of csi1p in other species, implying that csi1p may not be conserved through evolution or has evolved to functional homologues with a short conserved domain as alp7p/TACC (Peset and Vernos, 2008). Despite this, all csi1p binding proteins reported thus far, including sad1p, spc7p (Hou *et al.*, 2012), and alp7p (Figure 1), are well conserved through evolution. In addition, the mechanism targeting alp7p/TACC to the centrosome/SPB is conserved. Therefore, to identify csi1p homologues in other species, it will be of great interest to explore the centrosomal proteins that are involved in bipolar spindle formation and have the ability to interact with the conserved TACC domain.

MATERIALS AND METHODS

Plasmids and yeast strains

Yeast genetics was carried out as previously described (Forsburg and Rhind, 2006), and yeast strains were created by either random spore digestion or tetrad dissection. Yeast culture media were purchased from ForMedium (Norfolk, UK). Mutagenesis was performed with the QuikChange II XL Site-Directed Mutagenesis Kit (Agilent Technologies, Santa Clara, CA). MBC sensitivity assays were carried out on YES5S (yeast extract medium supplemented with adenine, leucine, uracil, histidine, and lysine) plates supplied with the indicated concentration of the microtubule-depolymerizing drug MBC (Sigma-Aldrich, St. Louis, MO). Minichromosome loss assays were carried out as described in Niwa *et al.* (1989). All yeast strains and plasmids used in this study are listed in Supplemental Tables S1 and S2, respectively.

Yeast two-hybrid assays

Yeast two-hybrid assays were performed using the Matchmaker Gold yeast two-hybrid system, along with Yeastmaker Yeast Transformation System 2 (Clontech, Mountain View, CA). Tenfold serial dilutions of Y2HGold cells containing bait and prey plasmids were cultured on double-dropout medium SD/-Leu/-Trp plates and quadruple-dropout medium SD/-Leu/-Trp/-Ade/-Ura plates containing 40 μg/ml X-α-Gal and 200 ng/ml Aureobasidin A at 30°C for 4 d.

Biochemistry

Recombinant proteins were purified from *Escherichia coli* using either glutathione Sepharose 4B resins (for GST-fused proteins; GE Healthcare, Pittsburgh, PA) or nickel resins (for His-tagged proteins; Qiagen, Valencia, CA). GST pull-down assays were then carried out by incubating GST-fused proteins bound to the glutathione resins with His-tagged proteins in Tris-buffered saline (TBS) plus 0.1% Triton X-100 at 4°C for 4 h, followed by 5× intensive washing with the TBS plus 0.1% Triton X-100 buffer and 1× TBS buffer. The resulting pull-down products were analyzed by silver staining and Western blotting with anti-His antibody (GE Healthcare). For protein expression level analysis, yeast protein extract was prepared as previously described (Matsuo *et al.*, 2006), followed by SDS-PAGE analysis and Western blotting with anti-GFP (Rockland Immunochemicals, Gilbertsville, PA) and anti-tubulin antibodies.

Microscopy and data analysis

Imaging was carried out as previously described (Tran *et al.*, 2004). Briefly, a PerkinElmer spinning-disk confocal microscope equipped with a Zeiss PlanApo 100×/1.4 numerical aperture objective and the Photometrics electron-multiplying charge-coupled device camera Evolve 512 was used to carry out live-cell imaging at 26°C in a temperature-controllable incubator. For maximum projection analysis, Z-stack images consisting of 11 planes (step size, 0.5 µm) were acquired every 30 s (for spindle dynamics analysis) or 1 min (for chromosome dynamics analysis); for sum projection analysis, Z-stack images consisting of 21 planes (step size, 0.25 µm) were acquired. Detailed imaging conditions are also described in the Supplemental Material. Spindle lengths were measured with MetaMorph (Molecular Devices, Sunnyvale, CA) and the MTrackJ plug-in in ImageJ (National Institutes of Health, Bethesda, MD) as previously described (Fu *et al.*, 2009). Fluorescence intensity measurements were performed using the Linescan function in MetaMorph, and values were then normalized to the maximum fluorescence intensity in each comparison group. Student's *t* tests were determined using Excel. Box plots and graphs were generated with Kaleidagraph 4.5 (Synergy Software, Reading, PA).

Note added in proof. While our manuscript was in press, Takashi Toda (Cancer Research UK) and colleagues published a paper showing that pcp1p is also involved in the recruitment of alp7p to the SPB (Tang *et al.*, 2014).

ACKNOWLEDGMENTS

We thank Takashi Toda (Cancer Research UK, London, United Kingdom), Snezhana Oliferenko (King's College London, London, United Kingdom), Dannel McCollum (University of Massachusetts Medical School, Worcester, MA), Yoshinori Watanabe (University of Tokyo, Japan), Mitsuhiro Yanagida (University of Tokyo, Japan), and Fred Chang (Columbia University, New York, NY) for providing yeast strains. We also thank George Tsao and Jing Guo of the Faculty Core Facility at the University of Hong Kong for technical support. This work is supported by grants from the National Natural Science Foundation of China (31271439 and 31301109), the Hong Kong Research Grants Council, and the Hong Kong University Seed Funding Programme to C.F., the S.K. Yee Medical Research Fund 2011, and grants from the National Institutes of Health and the Agence Nationale de la Recherche to P.T.T.

REFERENCES

Al-Bassam J, Kim H, Flor-Parra I, Lal N, Velji H, Chang F (2012). Fission yeast Alp14 is a dose-dependent plus end-tracking microtubule polymerase. *Mol Biol Cell* 23, 2878–2890.

Bellanger JM, Gonczy P (2003). TAC-1 and ZYG-9 form a complex that promotes microtubule assembly in *C. elegans* embryos. *Curr Biol* 13, 1488–1498.

Booth DG, Hood FE, Prior IA, Royle SJ (2011). A TACC3/ch-TOG/clathrin complex stabilises kinetochore fibres by inter-microtubule bridging. *EMBO J* 30, 906–919.

Costa J, Fu C, Syrovatkin V, Tran PT (2013). Imaging individual spindle microtubule dynamics in fission yeast. *Methods Cell Biol* 115, 385–394.

Forsburg SL, Rhind N (2006). Basic methods for fission yeast. *Yeast* 23, 173–183.

Fu C, Ward JJ, Loidice I, Velve-Casquillas G, Nedelec FJ, Tran PT (2009). Phospho-regulated interaction between kinesin-6 Klp9p and microtubule bundler Ase1p promotes spindle elongation. *Dev Cell* 17, 257–267.

Fu W, Tao W, Zheng P, Fu J, Bian M, Jiang Q, Clarke PR, Zhang C (2010). Clathrin recruits phosphorylated TACC3 to spindle poles for bipolar spindle assembly and chromosome alignment. *J Cell Sci* 123, 3645–3651.

Gergely F, Draviam VM, Raff JW (2003). The ch-TOG/XMAP215 protein is essential for spindle pole organization in human somatic cells. *Genes Dev* 17, 336–341.

Gergely F, Karlsson C, Still I, Cowell J, Kilmartin J, Raff JW (2000a). The TACC domain identifies a family of centrosomal proteins that can interact with microtubules. *Proc Natl Acad Sci USA* 97, 14352–14357.

Gergely F, Kidd D, Jeffers K, Wakefield JG, Raff JW (2000b). D-TACC: a novel centrosomal protein required for normal spindle function in the early *Drosophila* embryo. *EMBO J* 19, 241–252.

Giet R, McLean D, Descamps S, Lee MJ, Raff JW, Prigent C, Glover DM (2002). *Drosophila* Aurora A kinase is required to localize D-TACC to centrosomes and to regulate astral microtubules. *J Cell Biol* 156, 437–451.

Hiraoka Y, Toda T, Yanagida M (1984). The NDA3 gene of fission yeast encodes beta-tubulin: a cold-sensitive nda3 mutation reversibly blocks spindle formation and chromosome movement in mitosis. *Cell* 39, 349–358.

Hood FE, Williams SJ, Burgess SG, Richards MW, Roth D, Straube A, Pfuhl M, Bayliss R, Royle SJ (2013). Coordination of adjacent domains mediates TACC3-ch-TOG-clathrin assembly and mitotic spindle binding. *J Cell Biol* 202, 463–478.

Hou H, Zhou Z, Wang Y, Wang J, Kallgren SP, Kurchuk T, Miller EA, Chang F, Jia S (2012). Csi1 links centromeres to the nuclear envelope for centromere clustering. *J Cell Biol* 199, 735–744.

Janson ME, Loughlin R, Loidice I, Fu C, Brunner D, Nedelec FJ, Tran PT (2007). Crosslinkers and motors organize dynamic microtubules to form stable bipolar arrays in fission yeast. *Cell* 128, 357–368.

Kapitein LC, Peterman EJ, Kwok BH, Kim JH, Kapoor TM, Schmidt CF (2005). The bipolar mitotic kinesin Eg5 moves on both microtubules that it crosslinks. *Nature* 435, 114–118.

Kodama Y, Hu CD (2012). Bimolecular fluorescence complementation (BiFC): a 5-year update and future perspectives. *BioTechniques* 53, 285–298.

Le Bot N, Tsai MC, Andrews RK, Ahringer J (2003). TAC-1, a regulator of microtubule length in the *C. elegans* embryo. *Curr Biol* 13, 1499–1505.

Lin CH, Hu CK, Shih HM (2010). Clathrin heavy chain mediates TACC3 targeting to mitotic spindles to ensure spindle stability. *J Cell Biol* 189, 1097–1105.

Marguerat S, Schmidt A, Codlin S, Chen W, Aebersold R, Bahler J (2012). Quantitative analysis of fission yeast transcriptomes and proteomes in proliferating and quiescent cells. *Cell* 151, 671–683.

Matsuo Y, Asakawa K, Toda T, Katayama S (2006). A rapid method for protein extraction from fission yeast. *Biosci Biotechnol Biochem* 70, 1992–1994.

Mori D, Yano Y, Toyo-oka K, Yoshida N, Yamada M, Muramatsu M, Zhang D, Saya H, Toyoshima YY, Kinoshita K, *et al.* (2007). NDEL1 phosphorylation by Aurora-A kinase is essential for centrosomal maturation, separation, and TACC3 recruitment. *Mol Cell Biol* 27, 352–367.

Niwa O, Matsumoto T, Chikashige Y, Yanagida M (1989). Characterization of *Schizosaccharomyces pombe* minichromosome deletion derivatives and a functional allocation of their centromere. *EMBO J* 8, 3045–3052.

O'Brien LL, Albee AJ, Liu L, Tao W, Dobryzn P, Lizarraga SB, Wiese C (2005). The *Xenopus* TACC homologue, maskin, functions in mitotic spindle assembly. *Mol Biol Cell* 16, 2836–2847.

Peset I, Seiler J, Sardon T, Bejarano LA, Rybina S, Vernos I (2005). Function and regulation of maskin, a TACC family protein, in microtubule growth during mitosis. *J Cell Biol* 170, 1057–1066.

- Peset I, Vernos I (2008). The TACC proteins: TACC-ling microtubule dynamics and centrosome function. *Trends Cell Biol* 18, 379–388.
- Peterman EJ, Scholey JM (2009). Mitotic microtubule crosslinkers: insights from mechanistic studies. *Curr Biol* 19, R1089–1094.
- Petersen J, Paris J, Willer M, Philippe M, Hagan IM (2001). The *S. pombe* Aurora-related kinase Ark1 associates with mitotic structures in a stage dependent manner and is required for chromosome segregation. *J Cell Sci* 114, 4371–4384.
- Royle SJ (2012). The role of clathrin in mitotic spindle organisation. *J Cell Sci* 125, 19–28.
- Sato M, Toda T (2007). Alp7/TACC is a crucial target in Ran-GTPase-dependent spindle formation in fission yeast. *Nature* 447, 334–337.
- Sato M, Vardy L, Angel Garcia M, Koonrugsa N, Toda T (2004). Interdependency of fission yeast Alp14/TOG and coiled coil protein Alp7 in microtubule localization and bipolar spindle formation. *Mol Biol Cell* 15, 1609–1622.
- Silkworth WT, Cimini D (2012). Transient defects of mitotic spindle geometry and chromosome segregation errors. *Cell Div* 7, 19.
- Srayko M, Quintin S, Schwager A, Hyman AA (2003). *Caenorhabditis elegans* TAC-1 and ZYG-9 form a complex that is essential for long astral and spindle microtubules. *Curr Biol* 13, 1506–1511.
- Subramanian R, Ti SC, Tan L, Darst SA, Kapoor TM (2013). Marking and measuring single microtubules by PRC1 and kinesin-4. *Cell* 154, 377–390.
- Syrovatkina V, Fu C, Tran PT (2013). Antagonistic spindle motors and MAPs regulate metaphase spindle length and chromosome segregation. *Curr Biol* 23, 2423–2429.
- Tanenbaum ME, Medema RH (2010). Mechanisms of centrosome separation and bipolar spindle assembly. *Dev Cell* 19, 797–806.
- Tang NH, Okada N, Fong CS, Arai K, Sato M, Toda T (2014). Targeting Alp7/TACC to the spindle pole body is essential for mitotic spindle assembly in fission yeast. *FEBS Lett*, doi: 10.1016/j.febslet.2014.06.027.
- Tang NH, Takada H, Hsu KS, Toda T (2013). The internal loop of fission yeast Ndc80 binds Alp7/TACC-Alp14/TOG and ensures proper chromosome attachment. *Mol Biol Cell* 24, 1122–1133.
- Thadani R, Ling YC, Oliferenko S (2009). The fission yeast TACC protein Mia1p stabilizes microtubule arrays by length-independent crosslinking. *Curr Biol* 19, 1861–1868.
- Tran PT, Paoletti A, Chang F (2004). Imaging green fluorescent protein fusions in living fission yeast cells. *Methods* 33, 220–225.
- Usui T, Maekawa H, Pereira G, Schiebel E (2003). The XMAP215 homologue Stu2 at yeast spindle pole bodies regulates microtubule dynamics and anchorage. *EMBO J* 22, 4779–4793.
- Walczak CE, Heald R (2008). Mechanisms of mitotic spindle assembly and function. *Int Rev Cytol* 265, 111–158.
- Zhang J, Megraw TL (2007). Proper recruitment of gamma-tubulin and D-TACC/Msps to embryonic *Drosophila* centrosomes requires centrosomin motif 1. *Mol Biol Cell* 18, 4037–4049.